



(11) **EP 1 547 642 B1**

(12) **EUROPEAN PATENT SPECIFICATION**

(45) Date of publication and mention  
of the grant of the patent:  
**08.08.2007 Bulletin 2007/32**

(51) Int Cl.:  
**A61M 39/02 (2006.01)**

(21) Application number: **04257796.5**

(22) Date of filing: **15.12.2004**

(54) **Flexible injection port**

Flexible Injektionsöffnung

Orifice d'injection souple

(84) Designated Contracting States:  
**AT BE BG CH CY CZ DE DK EE ES FI FR GB GR  
HU IE IS IT LI LT LU MC NL PL PT RO SE SI SK TR**

(30) Priority: **16.12.2003 US 737942**

(43) Date of publication of application:  
**29.06.2005 Bulletin 2005/26**

(60) Divisional application:  
**06076702.7 / 1 733 756**

(73) Proprietor: **ETHICON ENDO-SURGERY, INC.**  
**Cincinnati, Ohio 45242 (US)**

(72) Inventors:  
• **Chen, How-Lun**  
**Cincinnati, Ohio 45242 (US)**

• **Conlon, Sean P.**  
**Loveland, Ohio 45140 (US)**  
• **Schulze, Dale R.**  
**Lebanon, Ohio 45036 (US)**

(74) Representative: **Tunstall, Christopher Stephen et  
al**  
**Carpmaels & Ransford**  
**43-45 Bloomsbury Square**  
**London WC1A 2RA (GB)**

(56) References cited:  
**EP-A- 0 203 633 WO-A-00/27299**  
**US-A- 4 160 454 US-A- 4 184 497**  
**US-A- 5 207 644**

**EP 1 547 642 B1**

Note: Within nine months from the publication of the mention of the grant of the European patent, any person may give notice to the European Patent Office of opposition to the European patent granted. Notice of opposition shall be filed in a written reasoned statement. It shall not be deemed to have been filed until the opposition fee has been paid. (Art. 99(1) European Patent Convention).

## Description

### Field of the Invention

[0001] This invention relates generally to the field of medicine, and more specifically to medical devices that are surgically implanted in a patient, and is particularly relevant to implantable injection or infusion ports such as used for chemotherapy and adjustable gastric band procedures.

### Background

[0002] Surgeons routinely implant subcutaneous injection ports in patients requiring long term, periodic fluid injections such as for chemotherapy and gastric band adjustments. The injection port connects to a flexible tube catheter to transport the fluid to the affected area (subclavian vein, etc.) or the gastric band. Current injection ports comprise a rigid metal or plastic housing, which is about 25mm in diameter and 15mm tall. A thick, silicone septum captured within the rigid housing covers an inner chamber that fluidly communicates with the catheter. The surgeon uses a hypodermic needle to inject fluid into the chamber through the silicone septum.

[0003] Typically the surgeon fastens the injection port with suture to fascia and beneath the fat and skin layers, primarily to prevent the port from flipping over, but also to prevent the injection port from migrating in the body. Since the septum is accessible from only one side of the injection port, flipping over requires interventional surgery to right the port for subsequent injections.

[0004] For some patients, the surgeon may place the injection port in the lower abdomen, thus burying the port beneath a fat layer that may be several centimeters thick. Usually a surgeon can locate the port with palpation alone. However, if there is a very thick, intervening fat layer, such as on extremely obese, gastric band patients, the surgeon must also use fluoroscopy, ultrasound, or other means to locate the port. Furthermore, the surgeon must inject the needle in a direction approximately perpendicular to the injection port, and hit the target area of the septum, which is only about 12-15mm in diameter. For some patients, the surgeon may place the injection port on the sternum or upper right chest, just beneath the skin layers. Although easy to locate with palpation, some patients regard the protruding port as uncomfortable or cosmetically objectionable.

[0005] EP-A-0203633 discloses an injection port for subcutaneous placement within a body. The injection port includes a rigid body having a membrane sealingly attached thereto. The membrane is such that it will self seal after being punctured by a needle. The body of the device and the membrane together define a fluid reservoir. The body and the membrane are both covered by a flexible silicone cover. The injection port includes a flexible elongated tubular catheter attached to the body which is in fluid communication with the reservoir.

[0006] US 4,160,454 and US 4,184,497 each disclose an implantable catheter system including a flexible catheter connected to a body. The body is an elongate tube and comprises a self sealing silicone wall defining a fluid reservoir within. The wall may include an embedded wire coil to provide support.

[0007] US 5,207,644 discloses an injection port for subcutaneous placement within a body. The injection port includes a body having a flexible catheter connected thereto. The body includes an expandable wall having an injection window formed of self sealing material. The wall and window together define a fluid reservoir.

[0008] WO 00/27299 discloses an implantable flexible tube having a flexible catheter attached thereto. The tube comprises a self sealing wall.

[0009] What is needed, therefore, is a subcutaneously implantable injection port that is made of relatively soft and flexible materials, and ideally, that looks and feels more (than current injection ports) like a large, natural blood vessel. What is also needed is a subcutaneously implantable injection port that is penetrable with a hypodermic needle, independent of the orientation of the injection port in bodily tissue, and that is self-sealing when the needle is removed. What is further needed is a subcutaneously implantable injection port that a surgeon may position in the body more quickly and with less dissection than is required for conventional injection ports.

### Summary of the Invention

[0010] An injection port for subcutaneous placement within a body. The injection port includes an elongated flexible substantially non-rigid body having first and second ends and a wall therebetween. The wall is such that it will self seal after being punctured by a needle. The body of the device further includes a fluid reservoir surrounded by the wall. Lastly, the injection port includes a flexible elongated tubular catheter attached to the body which is in fluid communication with the reservoir. The wall comprises an inner layer and an outer layer, wherein the outer layer is in compression around the inner layer, and an inner lining tightly assembled inside of the inner layer to provide support to the inner layer.

### Brief Description of the Drawings

[0011] We present the specific, novel features of this invention in the appended claims. The reader may best understand, however, the organization and methods of operation of this invention by referring to the detailed description and the following drawings:

FIG. 1 is an isometric view of an injection port of the prior art;

FIG. 2 is a cross sectional view of the injection port of the prior art shown in FIG. 1;

FIG. 3 is an isometric view of a first embodiment of a flexible injection port 30;

FIG. 4 is a sectional view of flexible injection port 30 shown in FIG. 3;

FIG. 5 is an enlarged, longitudinal sectional view of flexible injection port 30 penetrated by a hypodermic needle 100;

FIG. 6 is a cross sectional view of a second embodiment of a flexible injection port 50;

FIG. 7 is a cross sectional view of a third embodiment of a flexible injection port 60;

FIG. 8 is an isometric view of a fourth embodiment of a flexible injection port 80;

FIG. 9 is a cross sectional view of flexible injection port 80;

FIG. 10 shows injection port 30 subcutaneously implanted near a fascia layer 124 in a patient;

FIG. 11 shows injection port 30 subcutaneously implanted near a skin layer 120 in a patient; and

FIG. 12 shows injection port 30 subcutaneously implanted in a fat layer 122 in a patient.

#### Detailed Description of the Invention

[0012] Referring now to the drawings, FIGS. 1 and 2 show an injection port 10 of the prior art. Injection port 10 generally has a truncated, conical configuration, and comprises a body portion 12, a housing 14, a seal element 16, and a catheter element 18. The body portion 12 is made of a flexible, rubberized material with a cavity 20 formed inside. A catheter support 22 integrally forms in body portion 12. Housing 14 is made of a corrosion resistant metal, and has a reduced, upwardly facing entry passage 24. Seal element 16 is made of a rubberized material, which is easily penetrable by a hypodermic needle or the like, and provides a penetrable seal for passage 24. Housing 14 and seal element 16 define an open cavity 20 in injection port 10 for receiving and containing a fluid. Catheter element 18 extends through catheter support 22 of body portion 12 and through housing 14 so that catheter element 18 extends into cavity 20 for providing communication between cavity 20 and the exterior of injection port 10 for dispensing fluid from the cavity 20 into the body of a patient.

[0013] A surgeon implants injection port 10 subcutaneously in a patient. To introduce a fluid such as a medication or a saline solution, the surgeon inserts a hypodermic needle or the like into the patient so that the tip of the needle passes through seal element 16 and into

cavity 20. Due to the relatively small size of passage 24, each time the surgeon introduces a fluid into the patient, the surgeon must insert the needle through seal element 16 and the same localized area of the skin and tissue of the patient. Accordingly, seal element 16 may become significantly damaged and eventually develop a leak. Also, the localized skin area and underlying tissue may not heal in the desired manner. Further, because housing 14 is made of metal, it can cause barbing of the needle tip, causing increased trauma to the patient upon withdrawal of the needle. Still further, because of the truncated conical configuration of injection port 10 and the metallic construction of housing 14, injection port 10 can cause substantial discomfort to a patient, particularly if the area of the patient adjacent the injection port is accidentally bumped or bruised. In addition, because of the truncated conical configuration of injection port 10, it can cause a relatively unattractive mound on the body of a patient. Still further, since fluid can only be introduced in cavity 20 through passage 24, a surgeon must insert a needle into injection port 10 in substantially perpendicular relation to the skin so that often the adjacent area of tissue or skin of the patient cannot effectively support the needle.

[0014] When using injection port 10 of the prior art in a laparoscopic procedure such as implantation of a gastric band, it is necessary for the surgeon to assemble injection port 10 to catheter element 18 during the laparoscopic procedure. This is because injection port 10 is too large to pass through a standard size (12mm diameter) laparoscopic port, which is used for access to the stomach inside the abdominal cavity. The surgeon must introduce the gastric band and the catheter into the abdominal cavity without the injection port attached to the free end of the catheter. Once the surgeon has secured the gastric band around the stomach, the surgeon externalizes the free end of the catheter through the abdominal muscle and fascia layers, subcutaneous fat layer, and the skin to assemble the injection port to the free end of the catheter. Then the surgeon implants the injection port subcutaneously at the desired location on the patient's abdomen or chest. The surgeon must take extra time to assemble the injection port to the catheter. Also, the surgeon must skillfully connect the injection port to the catheter during less than ideal conditions. Consequently, there is the potential complication of an undiscovered leak developing at the connection of the catheter to the port.

[0015] FIG. 3 is an isometric view of a first embodiment, not having all of the features of the present invention, showing a flexible injection port or body 30, that generally comprises a first end 34, a second end 36, and a cylindrical injection portion 32 extending there between. A surgeon may use a hypodermic needle or the like to penetrate injection portion 32 and introduce a fluid such as a medication or saline solution into flexible injection port 30. Injection portion 32 self-seals when the surgeon removes the hypodermic needle. Injection portion 32 may

have a length, but is not limited to, approximately 5 - 20 cm. Injection portion 32 may have a diameter, but is not limited to, approximately 5 - 12 mm. A catheter 42 attaches to first end 34 and distributes fluid injected into flexible injection port 30 to another portion of the patient's body. Catheter 42 is made from a silicone rubber or other biocompatible polymer such as known in the art for application to conventional injection ports, such as shown in FIGs 1 and 2. A tether 38 having an eye loop 40 extends from second end 36. A surgeon may use a conventional surgical grasping instrument to grasp tether 38, or a surgical suture tied to eye loop 40, or a combination of both grasper and suture, to facilitate placement of flexible injection port 30 in the body.

**[0016]** Although flexible injection port 30 is shown in FIG. 3 to be essentially straight, it is possible to construct it with a curved or non-straight shape in order to facilitate placement in the body, or to conform to the body anatomy at the implant location. Since flexible injection port 30 is made of relatively soft and flexible materials, the surgeon may temporarily straighten it, for example, when introducing it into the body through a laparoscopic port.

**[0017]** FIG. 4 is a cross sectional view of flexible injection port 30, taken at line 4-4 of injection portion 32 as shown in FIG. 3. At this location and anywhere along the length of injection portion 32, flexible injection port 30 includes an outer tube 44 that may exert a radial, compressive force on an inner tube 46. Flexible injection port 30 includes a fluid reservoir 48 that extends the entire length of injection portion 32 and fluidly communicates with catheter 42. The total wall thickness is approximately in the range of 2 - 4 mm.

**[0018]** FIG. 5 is a longitudinal sectional view of flexible injection port 30, showing a hypodermic needle 100 penetrating through injection portion 32 so that distal tip 102 of hypodermic needle 100 is inside of fluid reservoir 48. First end 34, second end 36, tether 38, eye loop 40, and inner tube 46 are integrally molded from an elastomer such as, for example, silicone rubber, latex rubber, or polyurethane rubber. The molded elastomer may have a durometer approximately in the range of 40-60, but is not limited to that range. Catheter 42 may be bonded inside of first end 34 using any one of a number of bonding agents and techniques well known in the art, in order to fluidly communicate with reservoir 48. Outer tube 44 may be made of a PTFE shrink-wrap material, or a similar, biocompatible shrink-wrap. During the manufacturing process, outer tube 44 may be loosely assembled in the pre-shrunk configuration over inner tube 46. Then the application of heat causes outer tube 44 to conform very tightly around inner tube 46. Outer tube 44 therefore applies a significant compressive force on the softer, inner tube 46 to enhance the ability of inner tube 46 to close the puncture created by hypodermic needle 100.

**[0019]** FIG. 6 is a cross sectional view of a second embodiment comprising the present invention showing a flexible injection port 50, which is externally similar to the first embodiment shown in FIG. 3. Flexible injection

port 50 includes an outer tube 52, an inner tube 54, and an inner lining 56. Outer tube 52 and inner tube 54 are the same as outer tube 44 and inner tube 46, respectively, of the first embodiment in FIG. 4. Inner lining 56 may be an extruded plastic, thin wall tube, such as polyethylene or PTFE, tightly assembled inside of inner tube 54 to provide internal support to inner tube 54. By supporting inner tube 54 in this way, a greater compressive force may be applied by outer tube 52 onto inner tube 54, to further enhance the self-sealing capability. The material of inner lining 56 may be selected to have a higher needle penetration resistance than inner tube 54. This difference in penetration resistance provides the surgeon with tactile feedback that the needle tip has penetrated into fluid reservoir 58. Inner lining 56 may also be constructed of a metallic mesh and be similar in many respects to a vascular stent. Again, the total wall thickness is approximately in the range of 2 - 4 mm.

**[0020]** FIG. 7 is a cross sectional view of a third embodiment, not having all of the features of the present invention, showing a flexible injection port 60, which also is externally similar to the first embodiment shown in FIG. 3. Flexible injection port 60 comprises a plurality of layers 61, which for this third embodiment includes a first layer 62, a second layer 64, a third layer 66, a fourth layer 68, and a fifth layer 70, which surrounds a fluid reservoir 72.. Once penetrated by a needle that is inserted at an acute angle, the punctures created through the layers are not aligned to allow leakage once the needle is removed. Each of layers 61 may be made of the same or a different material than any of the other of layers 61, or may have the same or a different thickness than any of the other of layers 61. Each of layers 61 may have a specific property or functional contribution. For example, first layer 62 may be made of a material that is permeable to tissue fluids in order to slowly release a medication contained in second layer 64. Fifth layer 70 may be made of silicone rubber having a durometer in the range of 20 - 30. Fourth layer 68 may be made of a heat shrinkable PTFE material, which applies a radially compressive force on fifth layer 70 to enhance self-sealing. Third layer 66 may be made of a material such as a metallic foil that acts as a diffusion barrier to prevent the loss of fluid from fluid reservoir 72. Fourth layer 66 may be made of a high durometer silicone rubber. Many other materials are possible, in a multiplicity of combinations, so that injection port 60 may have characteristics especially suited for its particular application. Diffusion of body fluids into and out of the soft port wall may also be reduced by any one of various material treatment techniques, including, for example, vapor deposition of titanium or another metal on a surface of the soft port, and coating with *Paralene* polymer. Other coatings are also known in the art for micro bacterial protection. Again, the total wall thickness is in the range of 2 - 4 mm.

**[0021]** FIG. 8 is a fourth embodiment, not having all of the features of the present invention, a flexible injection port 80, comprising a first end 84 that attaches to a cath-

eter 92, a second end 86 and an injection portion 82. Flexible injection port 80 further comprises a webbing 88 attached to and covering at least injection portion 82, and made of a thin, flexible, implantable material such as a polyester or polypropylene mesh, expanded PTFE, or the like. Webbing 88 provides broad margins for stapling or suturing to an underlying tissue such as fascia, as well as a large area for tissue in-growth, to enhance long-term stability and to substantially prevent migration of flexible injection port 80. FIG. 9 is a cross sectional view of flexible injection port 80, taken at line 9-9 of FIG. 8. Flexible injection port 80 comprises an outer tube 94 made of a heat shrinkable, PTFE material, and an inner tube 96 made of a silicone rubber having a durometer of approximately 20-40. Webbing 88 includes a pair of webbing layers, 91 and 93, that may be bonded thermally or chemically tightly over at least injection portion 82 in the mid-plane of flexible injection port 80.

**[0022]** A surgeon may implant the present invention, as described for the preceding embodiments and equivalents, in a number of locations in a patient's body. FIGs 10, 11, and 12 show examples of flexible injection port 30 subcutaneously implanted in the abdomen of a patient, although it is possible to implant flexible injection port 30 beneath the skin in other portions of the body.

**[0023]** FIG. 10 depicts a first example of flexible injection port 30 subcutaneously implanted in a patient's body. Flexible injection port 30 lies adjacent to a fascia layer 124 covering an abdominal wall 126. Catheter 42 passes from the abdominal cavity 128 through an abdominal opening 132, which the surgeon used together with a first incision 130 for laparoscopic access earlier in the surgical procedure. The surgeon optionally may make a second incision 134 offset from first incision 130, and use conventional, surgical grasping and retracting instruments to pull flexible injection port 30 beneath a fat layer 122 and adjacent to fascia layer 124. However, the surgeon may determine that it is not necessary to make a second incision 134, and instead use first incision 130 to push flexible injection port 30 into position. In either situation, the surgeon dissects as little tissue as practical in order to save surgery time and to minimize the size of enclosed cavities that may collect tissue fluids and become sites for infection. The surgeon optionally may anchor flexible injection port 30 to fascia layer 124 with a stay suture 102. Once the surgeon has placed flexible injection port 30 in the desired location, the surgeon closes first incision 130 and second incision 134 using conventional sutures or staples.

**[0024]** FIG. 11 shows a second example of flexible injection port 30 subcutaneously implanted in a patient's body. Flexible injection port 30 lies immediately beneath skin layer 120 and above fat layer 122. Catheter 42 passes through first incision 130 and abdominal opening 132 (the original laparoscopic port site) into abdominal cavity 128. The surgeon may use finger or instrument dissection through first incision 130 to create a space under skin layer 120 for flexible injection port 30. The surgeon closes

first incision 130 using conventional sutures or staples. Normally it would not be necessary to close abdominal opening 132 through fascia layer 124 and abdominal wall 126, but the surgeon may do so in order to promote healing and to prevent slippage of catheter 42 through abdominal opening 132. The surgeon may prefer placement of flexible injection port 30 just beneath skin layer 120 for severely obese patients in which fat layer 122 is over 5-10 cm thick, so that the surgeon may easily use palpation to locate flexible injection port 30 for later injections of fluid. Also, conventional intravenous (IV) needles and techniques may be used for injecting the fluid into flexible injection port 30, which is situated beneath the skin much like a natural blood vessel. This may allow nurses and other clinicians who are trained in administering IV's to assist the surgeon with fluid injections. Furthermore, if the clinician uses a conventional IV needle, the "flash-back" of fluid into the IV needle syringe tip provides the clinician with visual feedback that the tip of the needle is properly penetrated into the reservoir of flexible injection port 30. In fact, addition of a colorant to the fluid injected further enhances this visual feedback. Non-toxic colorants that may be added to the saline solution or medication are well known in the art.

**[0025]** FIG. 12 shows a third example of flexible injection port 30 subcutaneously implanted in a patient's body. For this example, the surgeon does minimal or no dissection of tissue at the laparoscopic port site. Catheter 42 passes from the abdominal cavity 128 through fascia layer 124 and abdominal wall 126. The surgeon positions flexible injection port 30 vertically in fat layer 122 and beneath skin layer 120. Optionally, the surgeon may suture abdominal opening 132 to prevent slippage of flexible injection port 30 into abdominal cavity 128. The surgeon also may use a surgical scissors to trim off tether 38 from flexible injection port 30, just prior to closing first incision 130 with conventional sutures or staples.

**[0026]** The present invention, a flexible injection port, as described in the preceding embodiments and their equivalents, has numerous advantages over the prior art injection ports. The flexible injection port may not require attachment to fascia, thus reducing the duration of the surgical procedure. The flexible injection port may require a smaller incision size and less tissue dissection for implantation, so that the patient has less pain, less scarring, a faster recovery, and less possibility of infection. Due to the integral construction of the flexible injection port and the catheter, the step of connecting the catheter to the injection port during the surgical procedure is not necessary, thus potentially reducing the number of surgical complications due to fluid leakage at the connection. Because the flexible injection port may be implanted in the fat layer near the skin surface, the surgeon or a trained clinician may use palpation to locate the injection port, and standard IV techniques to administer fluid, yet the implant is still cosmetically acceptable to the patient. In addition, shorter injection needles may be used to reduce patient anxiety during fluid administration. The flexible

injection port may have no metallic parts, resulting in a flexible and lightweight implant for greater patient comfort and compatibility with magnetic resonance and fluoroscopic x-ray imaging. Finally, the injection portion of the flexible injection port is accessible with a hypodermic needle for most of the possible orientations of the flexible injection port within the subcutaneous fat layer of the patient.

**[0027]** While preferred embodiments of the present invention have been shown and described herein, it will be obvious to those skilled in the art that such embodiments are provided by way of example only. Numerous variations, changes, and substitutions will now occur to those skilled in the art without departing from the invention. For example, the injection port may be coated with an antimicrobial coating such as triclosan. For example, as would be apparent to those skilled in the art, the disclosures herein have equal application in robotic-assisted surgery. In addition, it should be understood that every structure described above has a function and such structure can be referred to as a means for performing that function. Accordingly, it is intended that the invention be limited only by the scope of the appended claims.

#### Claims

1. An injection port (50) for subcutaneous placement within a body comprising:

- a. an elongated flexible substantially non-rigid body having first and second ends and a wall therebetween, said wall is such that it will self seal after being punctured, said body further including a fluid reservoir (58) surrounded by said wall; and
- b. a flexible elongated tubular catheter (42) attached to said body which is in fluid communication with said reservoir;

wherein said wall comprises an inner layer (54) and an outer layer (52), wherein said outer layer is in compression around said inner layer, and an inner lining (56) tightly assembled inside of the inner layer to provide internal support to the inner layer.

2. The flexible injection port of Claim 1, wherein each of said inner layer (54) and said outer layer (52) are made from a polymer material.
3. The flexible injection port of Claim 2 wherein said outer layer (52) is made of a heat shrinkable material, and said inner layer (54) is made of an elastomer.
4. The flexible injection port of claim 1 wherein said wall comprises a layer (66) of fluid diffusion barrier material.

5. The flexible injection port of claim 1 further comprising a webbing (88) attached to and extending from an outer surface of said body.

6. The flexible injection port of Claim 1, wherein said body has a diameter no greater than 12mm.

7. The flexible injection port of any preceding claim, wherein said inner lining (56) is made of a polymer or a metallic mesh material having a higher puncture resistance than the inner layer (54) and outer layer (52).

8. The flexible injection port of Claim 7, wherein said polymer comprises polyethylene or PTFE.

9. The flexible injection port of any preceding claim, wherein the substantially non-rigid body is tubular.

10. The flexible injection port of claim 11, wherein said body has a length between said ends of about 5mm to about 20mm and an outer diameter from about 5mm to about 12mm.

#### Patentansprüche

1. Injektionsanschluß (50) zur subkutanen Anordnung in einem Körper, umfassend:

- a. einen länglichen, flexiblen, im wesentlichen nicht starren Grundkörper mit einem ersten und einem zweiten Ende und einer dazwischen befindlichen Wand, wobei die genannte Wand so ist, daß sie sich selbsttätig abdichtet, nachdem sie durchlöchert worden ist, wobei der genannte Grundkörper weiterhin einen Fluidvorrat (58) aufweist, der von der genannten Wand umgeben ist; und
- b. einen flexiblen, länglichen, rohrförmigen Katheter (42), der an dem genannten Grundkörper befestigt ist, und der sich in Fluidverbindung mit dem genannten Vorrat befindet;

wobei die genannte Wand eine innere Schicht (54) und eine äußere Schicht (52) aufweist, wobei sich die genannte äußere Schicht in einem komprimierten Zustand um die genannte innere Schicht herum befindet, und eine innere Auskleidung (56), die eng innerhalb der inneren Schicht angeordnet ist, um eine innere Abstützung für die innere Schicht zu bilden.

2. Flexibler Injektionsanschluß nach Anspruch 1, dadurch gekennzeichnet, daß sowohl die genannte innere Schicht (54) als auch die genannte äußere Schicht (52) aus einem Polymermaterial hergestellt sind.

3. Flexibler Injektionsanschluß nach Anspruch 2, **dadurch gekennzeichnet, daß** die genannte äußere Schicht (52) aus einem heischrumpfbaren Material gebildet ist, und da die genannte innere Schicht (54) aus einem Elastomer gebildet ist.

5

4. Flexibler Injektionsanschluß nach Anspruch 1, **dadurch gekennzeichnet, da** die genannte Wand eine Schicht (66) aus einem Fluiddiffusions-Sperrmaterial aufweist.

10

5. Flexibler Injektionsanschluß nach Anspruch 1, weiter umfassend ein Gewebe (88), das an einer äußeren Oberfläche des genannten Grundkörpers befestigt ist und sich von dieser erstreckt.

15

6. Flexibler Injektionsanschluß nach Anspruch 1, **dadurch gekennzeichnet, da** der genannte Grundkörper einen Durchmesser aufweist, der nicht größer als 12 mm ist.

20

7. Flexibler Injektionsanschluß nach einem der vorangehenden Ansprüche, **dadurch gekennzeichnet, da** die genannte innere Auskleidung (56) aus einem Polymer oder aus einem metallischen Netzmaterial besteht, welches eine größere Widerstandsfähigkeit gegenüber Durchlöcherung aufweist als die innere Schicht (54) und die äußere Schicht (52).

25

8. Flexibler Injektionsanschluß nach Anspruch 7, **dadurch gekennzeichnet, da** das genannte Polymer Polyethylen oder PTFE umfat.

30

9. Flexibler Injektionsanschluß nach einem der vorangehenden Ansprüche, **dadurch gekennzeichnet, da** der im wesentlichen nicht starre Grundkörper rohrförmig ist.

35

10. Flexibler Injektionsanschluß nach Anspruch 11, **dadurch gekennzeichnet, da** der genannte Grundkörper eine Länge zwischen den genannten Enden von etwa 5 mm bis etwa 20 mm und einen äußeren Durchmesser von etwa 5 mm bis etwa 12 mm aufweist.

40

45

## Revendications

1. Orifice d'injection (50) pour un placement sous-cutané dans un corps, comprenant :

50

- a. un corps allongé, souple, sensiblement non rigide ayant des première et seconde extrémités et une paroi entre celles-ci, ladite paroi est telle qu'elle se fermera d'elle-même après avoir été perforée, ledit corps comprenant en outre un réservoir de fluide (58) entouré par ladite paroi ; et
- b. un cathéter souple allongé tubulaire (42) fixé

55

au dit corps qui est en communication fluide avec ledit réservoir ;

dans lequel ladite paroi comprend une couche intérieure (54) et une couche extérieure (52), dans lequel ladite couche extérieure est en compression autour de ladite couche intérieure, et un revêtement intérieur (56) est assemblé de manière serrée à l'intérieur de la couche intérieure pour fournir un support interne à la couche intérieure.

2. Orifice d'injection souple selon la revendication 1, dans lequel chacune de ladite couche intérieure (54) et de ladite couche extérieure (52) est faite à partir d'un matériau polymère.

3. Orifice d'injection souple selon la revendication 2, dans lequel ladite couche extérieure (52) est faite d'un matériau rétrécissant sous l'effet de la chaleur, et ladite couche intérieure (54) est faite d'un élastomère.

4. Orifice d'injection souple selon la revendication 1, dans lequel ladite paroi comprend une couche (66) d'un matériau faisant barrière à la diffusion de fluide.

5. Orifice d'injection souple selon la revendication 1, comprenant en outre une armature (88) fixée à et s'étendant à partir d'une surface extérieure dudit corps.

6. Orifice d'injection souple selon la revendication 1, dans lequel ledit corps a un diamètre qui n'est pas supérieur à 12 mm.

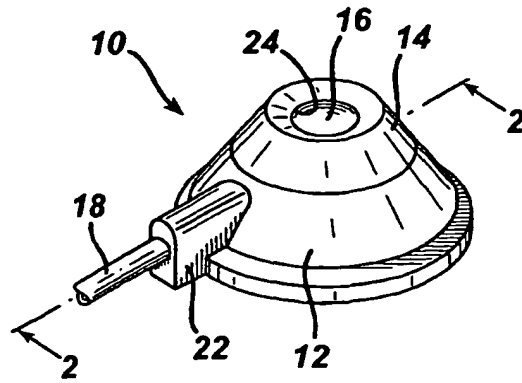
7. Orifice d'injection souple selon l'une quelconque des revendications précédentes, dans lequel ledit revêtement intérieur (56) est fait d'un polymère ou d'un matériau tressé métallique ayant une plus grande résistance à la perforation que la couche intérieure (54) et la couche extérieure (52).

8. Orifice d'injection souple selon la revendication 7, dans lequel ledit polymère comprend du polyéthylène ou du PTFE.

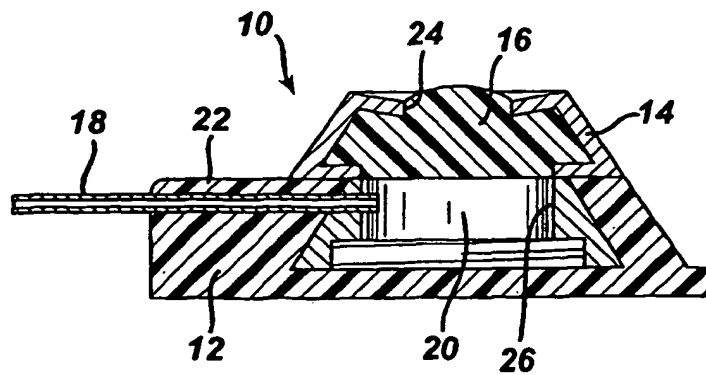
9. Orifice d'injection souple selon l'une quelconque des revendications précédentes, dans lequel ledit corps sensiblement non rigide est tubulaire.

10. Orifice d'injection souple selon la revendication 11, dans lequel ledit corps a une longueur entre lesdites extrémités d'environ 5 mm à environ 20 mm, et un diamètre extérieur d'environ 5 mm à environ 12 mm.

**FIG. 1** PRIOR ART



**FIG. 2** PRIOR ART





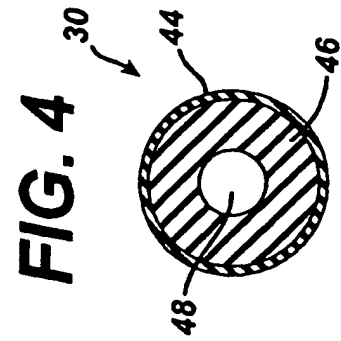
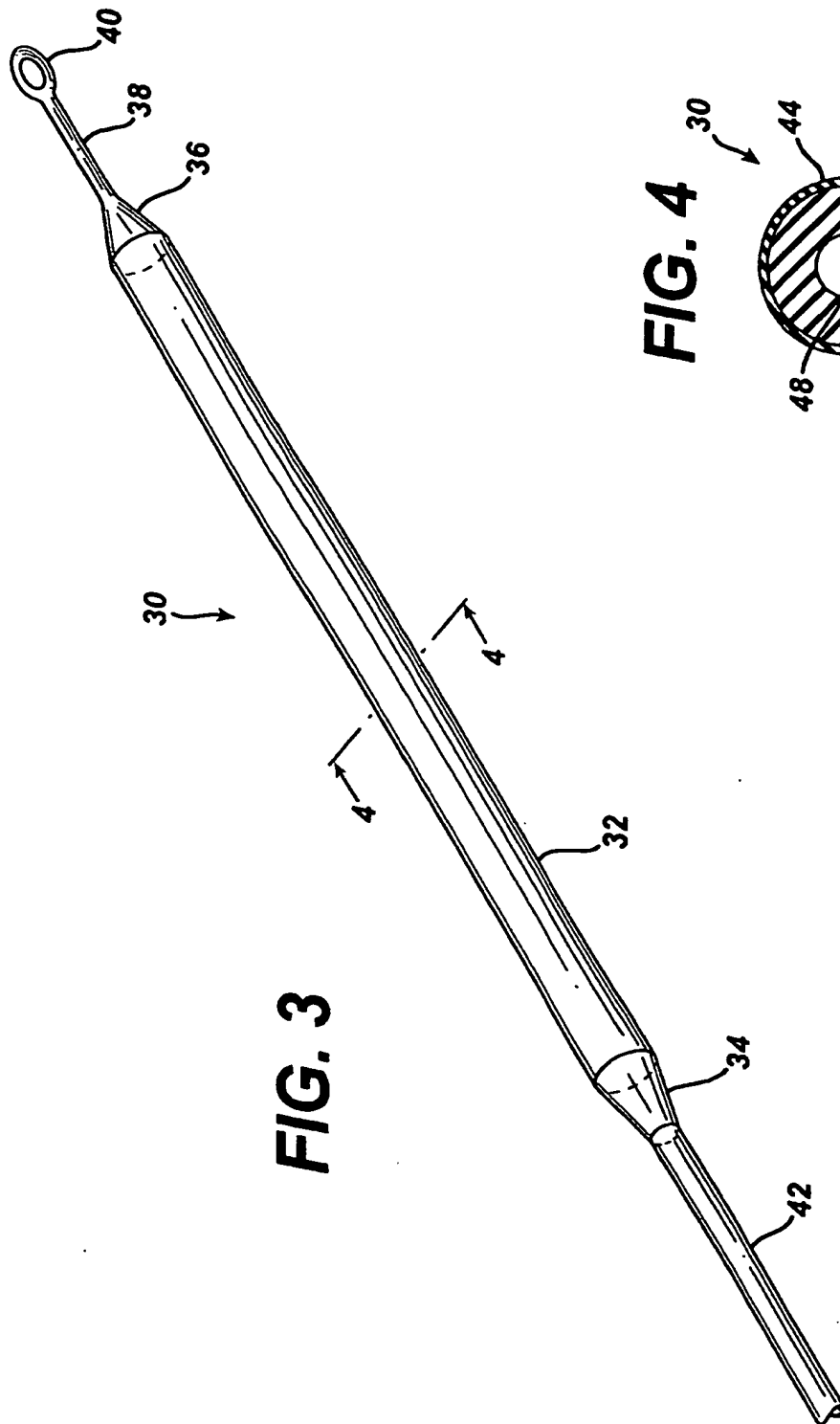
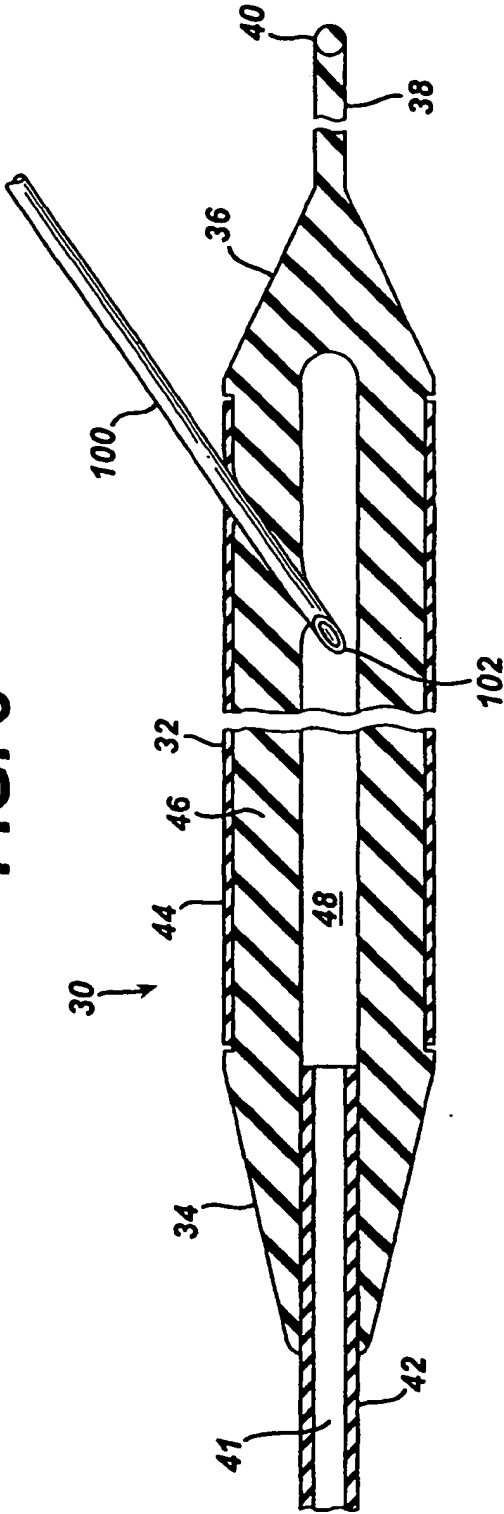
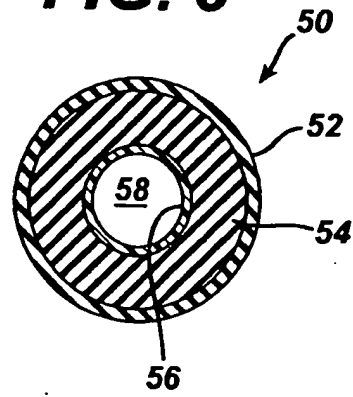


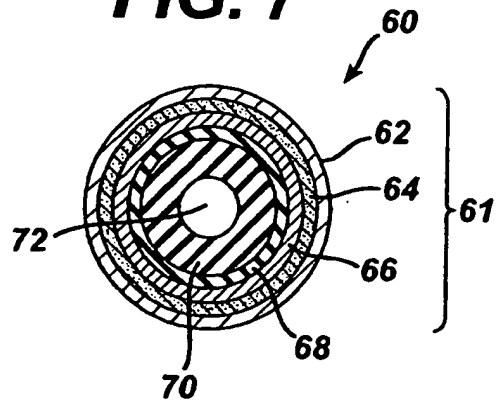
FIG. 5

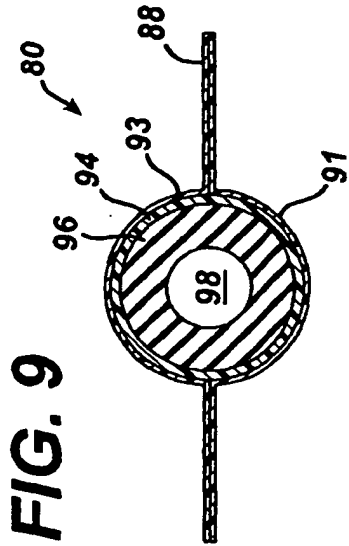
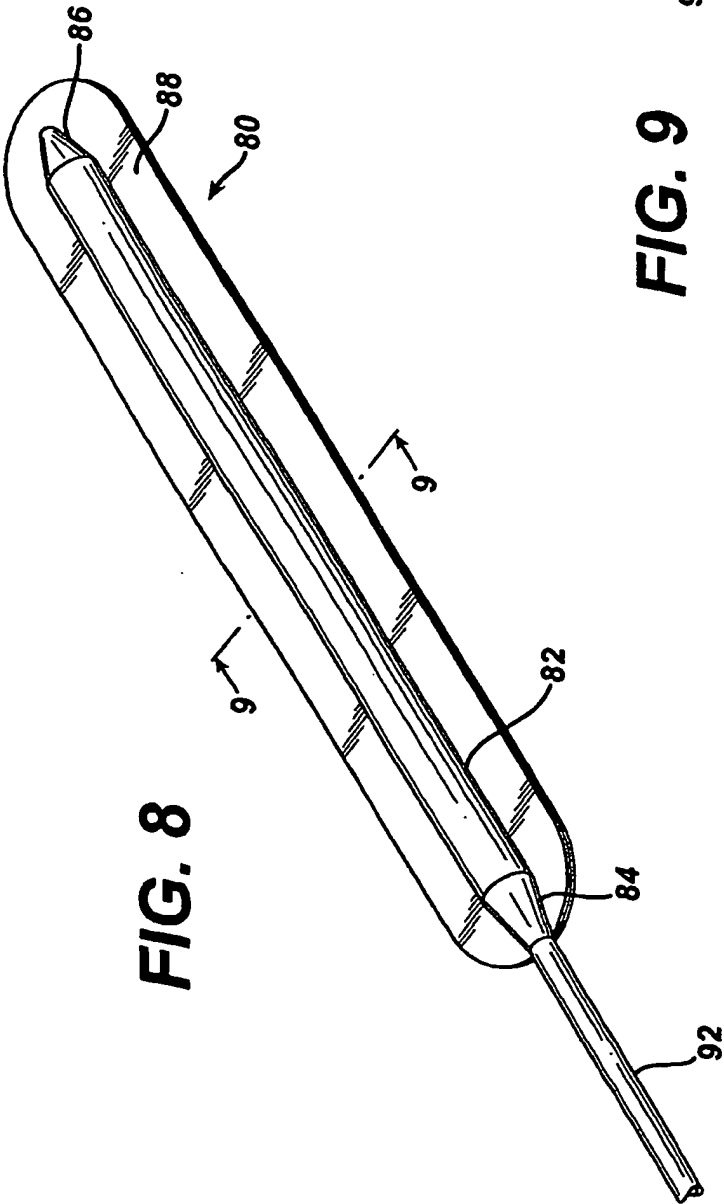


**FIG. 6**



**FIG. 7**





**FIG. 10**

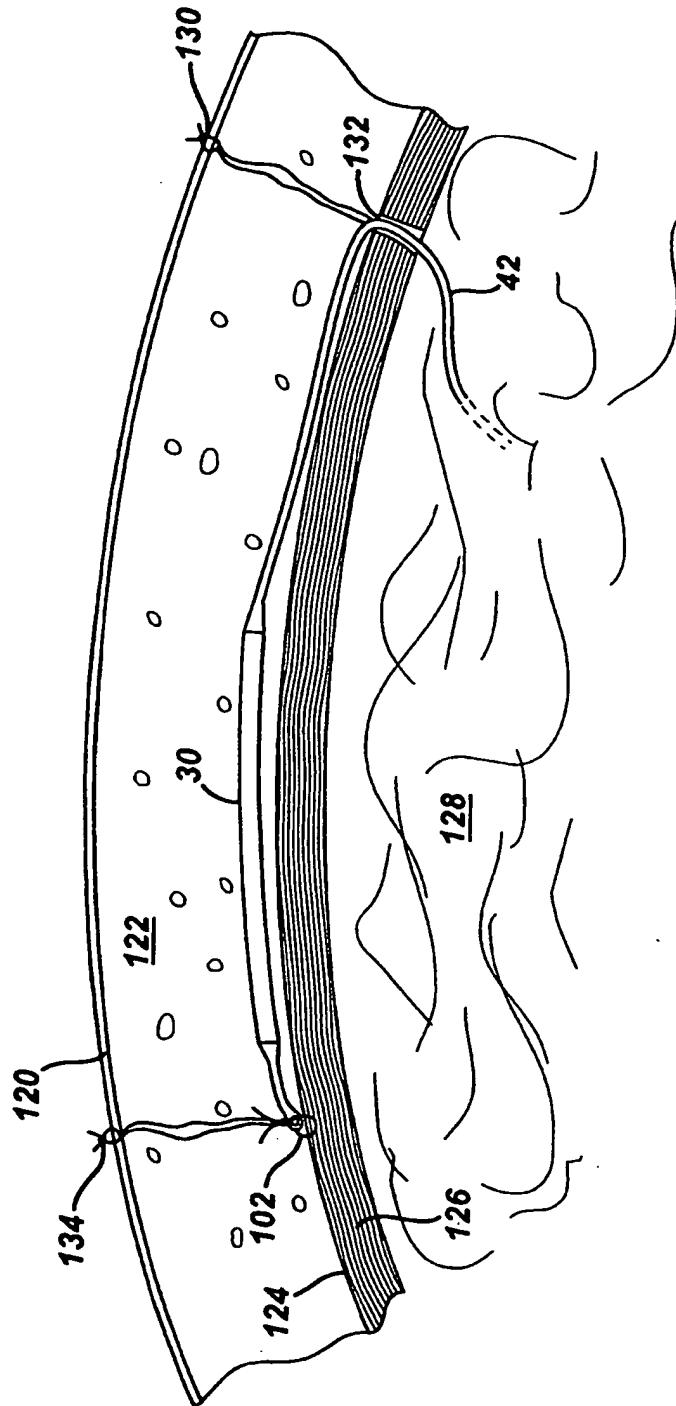
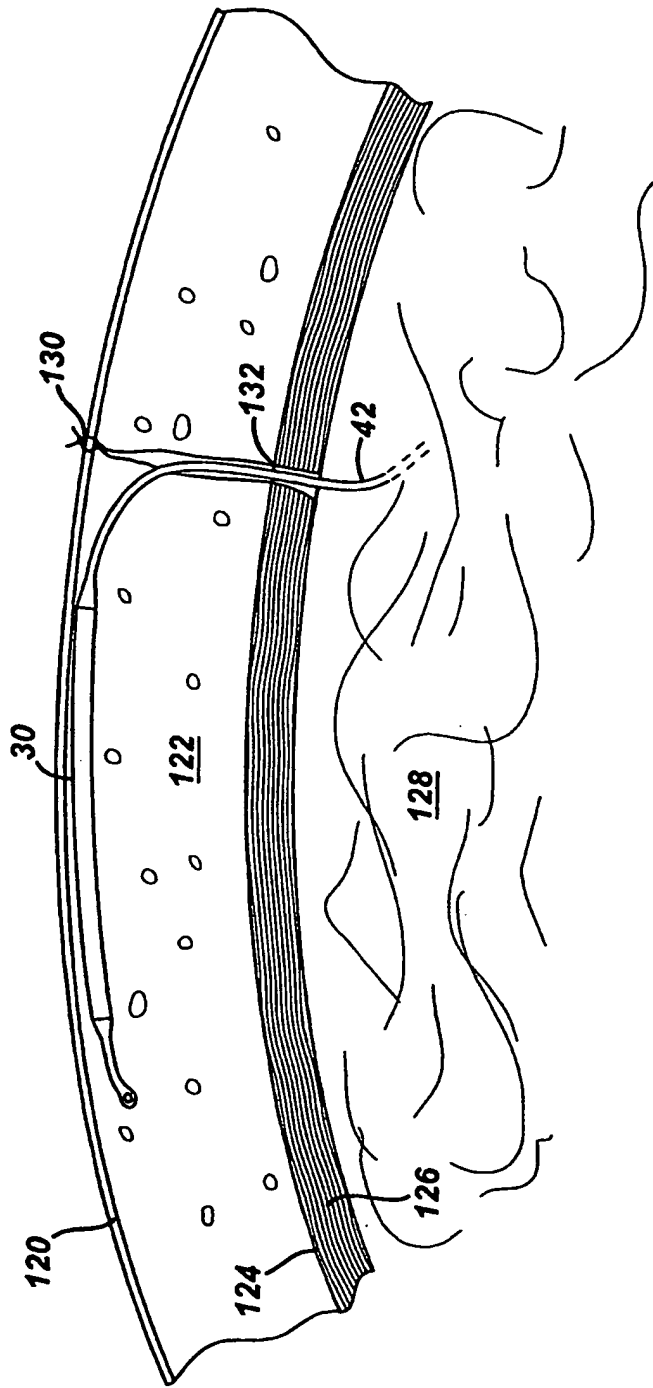
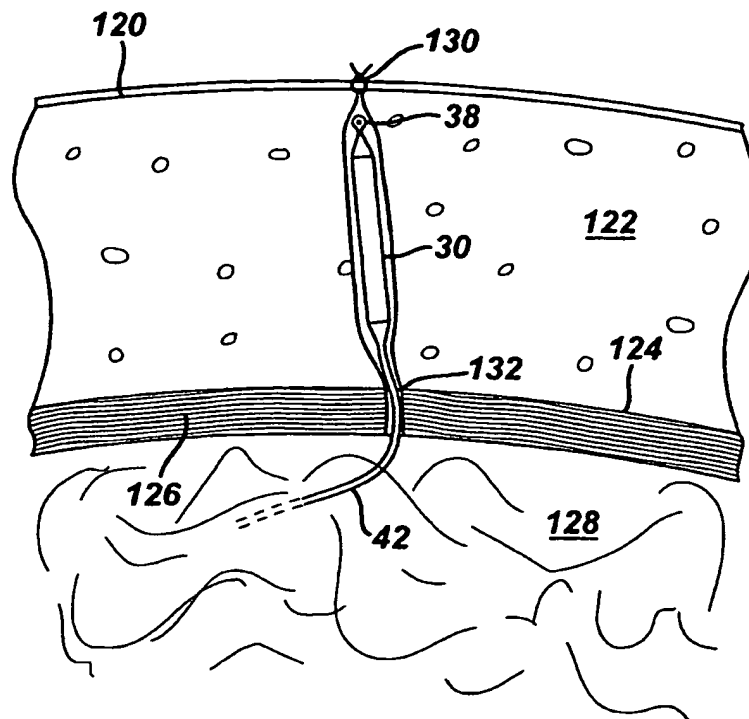


FIG. 11



**FIG. 12**



**REFERENCES CITED IN THE DESCRIPTION**

*This list of references cited by the applicant is for the reader's convenience only. It does not form part of the European patent document. Even though great care has been taken in compiling the references, errors or omissions cannot be excluded and the EPO disclaims all liability in this regard.*

**Patent documents cited in the description**

- EP 0203633 A [0005]
- US 4160454 A [0006]
- US 4184497 A [0006]
- US 5207644 A [0007]
- WO 0027299 A [0008]